

The **IREx Study Manager** is someone from the lead study team or coordinating center who uses IREx to oversee participating site readiness for single IRB (sIRB) review. For more detailed information on how to use IREx, check out the Study Manager Resources page [here](#).

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## Conversation with the Single IRB

- A. **Discuss** the submission process with the sIRB.
  - Review the [IREx Single IRB Instructions Template](#) for participating sites
  - Process for managing consent form (e.g., whether a template is being used)
  - Process for capturing local considerations from sites (e.g., via IREx surveys, including if the study will request local considerations updates throughout the life of the study)
  - Process for submitting sites for review (e.g., as an amendment, as a site add)
- B. **Clarify** what roles you are responsible for in IREx as a Study Manager vs the sIRB. Who will:
  - Upload Initial Approval for the Overall Study/Lead Site & Publish Approval
  - Add or remove participating sites to the study
  - Request Required Agreements
  - Grant study access to participating HRPP and study teams
  - Export participating sites' reliance and local review documentation for submission to the sIRB
  - Upload Reviewing IRB Site Approvals
  - Manage study-wide amendments and continuing reviews

**STEP****1****SUBMIT THE LEAD SITE TO THE SIRB**

- The **sIRB** will review the submission and create the study in IREx.
- You will receive access to IREx via an email notification after the study is created.
- While you wait for the sIRB to approve the Lead Site, you can complete steps 2 & 3.

**STEP****2****ADD PARTICIPATING SITES TO THE STUDY IN IREX****GETTING STARTED**

Upload Overall Study Approval

Publish Approval

**Add Participating Sites**

Request Agreements

Grant Site Access

View Site Progress on Status Summary

Upload Relying Site Approval

- A. Click **Add Participating Sites** in your Getting Started Checklist, this will open your **Sites** page.
- B. Search and add site(s) by the institutions' name (avoid abbreviations, e.g. "VUMC") or by Federalwide Assurance (FWA) # (numeric characters only) or select a consortium of sites.
- As you type, the sites that match the name/FWA # will appear. Select the site from drop-down list.
  - You can add sites that do not appear in the drop-down list by typing the site name/FWA # and pressing enter on your keyboard. An IREx admin will create the site in IREx and notify you when you can add contacts.
- C. Enter the PI and Study Team Member contact information, if known at the time. If not, **Save** the site and return later to add this information.

Approvals Status Summary Sites Contacts

Sites

Return to participating sites list

Add a participating site

Site

Island University - FWA0029993

List study team contacts that will need access to IREx. Go to Status Summary to grant study access.

☐ This Study Team engages additional sites

Personnel

Role	Email	First name	Last name
PI			
Study Team Member			

Add contact

Cancel Save

**STEP****3****REQUEST REQUIRED AGREEMENTS (IF AVAILABLE)**

- A. On the **Status Summary** page, any institution that is missing agreements will have an orange number of **Agreements Complete** button with a drop-down list of agreement(s) required for the study.
- B. Click the purple **Request Agreement(s)** button to send an email notification to the site.

Approvals Status Summary Sites Contacts

Status Summary

Export Data

Search:

Site	Agreements	Reliance Decision	Local Considerations	Approval Status
Carnegie University	2 / 2 Agreements Complete	Notify & Grant Access		
Evan University Medical Center	0 / 2 Agreements Complete	Notify & Grant Access		
Mellon University Medical Center	Reliance Agreement IREx Access	Notify & Grant Access		

Request Agreement(s)

## STEP

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## UPLOAD INITIAL SIRB APPROVAL FOR THE LEAD SITE/OVERALL STUDY & PUBLISH

GETTING STARTED

- Upload Overall Study Approval
- Publish Approval
- ✓ Add Participating Sites
- Grant Site Access
- View Site Progress on Status Summary
- Upload Relying Site Approval

- NOTE:** If applicable, include a site-specific consent template or part 2 consent template in the global documents.

A. Click **Upload Overall Study Approval** in your Getting Started Checklist.

B. Change the Status to **Approved** and indicate the reviewing type & cycle. All required fields and documents will be highlighted in red.

C. Enter the dates for when it was submitted, approved, review, and expires.

D. Upload your documents.

- Remember to click **Accept Draft** if the original protocol uploaded to IREx was approved or click **Replace Draft** if it was modified during the review, in which case be sure to change the version date BEFORE uploading the new protocol file.
- Replace or accept other draft documents, as needed.

Uploaded Documents

Review cannot be approved while documents are still in draft.

Type	Document	Date Approved
Protocol [1, 5/4/2023]	PROTOCOL_v1.docx DRAFT	

Accept Draft Replace Draft

- E. **Publish Approval** to make the approved global documents visible to the participating sites once they have study access. Check the box at the bottom of the Review & Submit page and click **Save**. Leave the box unselected and click **Save** if you want to return to Publish Approval later.

☒ Publish Lead Site / Overall study approval documents. If applicable, sites are not notified and cannot view these documents until their site approval is uploaded and saved.

Cancel Save

## STEP

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## NOTIFY & GRANT ACCESS TO SITES

A. Click **Grant Site Access** on your Getting Started Checklist, this will open your **Status Summary** page.

B. Click the **Notify & Grant Access** button to alert sites of their access to the study in IREx. This sends an email to the HRPP and study team to prompt them to connect around their local reliance process, gives them study access, and includes single IRB instructions for the study. The study team can use the Study Link in the email to log into IREx and download the lead site sIRB approval documents for their local submission (if applicable).

- Only sites that have joined IREx can be notified.
- You can notify sites at different times depending on when they are being onboarded to the study.

VUMC GETTING STARTED

- ✓ Add Participating Sites
- Grant Site Access
- View Site Progress on Status Summary
- Upload Relying Site Approval

Approvals Status Summary Sites Contacts Edit Study Info

### Status Summary

Export Data

Site	Agreements	Reliance Decision	Local Considerations	Approval Status
Carnegie University	2 / 2 Agreements Complete	Started 5/9/2023	0 / 3 Surveys Complete	Not Approved
Evan University Medical Center	0 / 2 Agreements Complete	Notify & Grant Access		
Mellon University Medical Center	2 / 2 Agreements Complete	Notify & Grant Access		

## STEP 6 TRACK SITES' READINESS FOR SIRB REVIEW

6 Use your **Status Summary** page to track your sites' progress.

**Status Summary**

Manage Agreements Export Data

Search:

Site	Agreements	Reliance Decision	Local Considerations	Approval Status
Anderson Medical Center	1 / 3 Agreements Complete	Notify & Grant Access		
Central Ohio Medical Center	3 / 3 Agreements Complete	Notify & Grant Access		
Faraday Institute of Research, Science and Technology	3 / 3 Agreements Complete	Contacted 4/13/2023		
Mellon University Medical Center	3 / 3 Agreements Complete	Completed 4/21/2023	3 / 3 Surveys Complete	Not Approved
Middle-earth College of Agriculture	3 / 3 Agreements Complete	Started 4/21/2023	1 / 3 Surveys Complete	Not Approved

Local Considerations Legend:

- ✓ Institutional Profile Confirmed: 4/21/2023
- ✗ HRP Survey
- ✗ PI Survey
- Email study personnel

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### A. Has the site signed all the required agreements?

- SMART IRB Agreement, IAA or MOU
- Indemnification (if applicable)
- IREx Access

### B. Has the site's HRPP indicated reliance?

- See Reliance Decision Legend – Gray is Completed

### C. Are local considerations complete?

- Institutional Profile: Completed by the HRPP and includes institution-level information.
- HRP Survey: Completed by the HRPP and includes applicable local requirements for this study.
- PI Survey: Completed by the PI or Study Team Member; includes information about the conduct of the study and an upload of the locally reviewed consent document(s). The PI must attest to the survey, as well as any edits made by a Study Team Member or the HRPP.

**IREx Tip:** If steps are incomplete, ensure the study team has submitted to their local IRB. If so, ask the study team to follow up with their IREx HRPP Liaison ([found here](#)) regarding steps in IREx or additional requirements.

Reliance Decision Legend	
Awaiting Confirmation	Reviewing IRB must accept SSRP
Add PI Info	Required PI information is not yet entered
Notify & Grant Access	Site has NOT joined IREx so cannot be granted access
Notify & Grant Access	Site has IREx access and can be notified of study access
Contacted	Site has access; click to re-send notification
Started	HRPP has accessed study in IREx
Completed	HRPP has ceded review

## STEP

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## PRE-SCREEN AND EXPORT LOCAL CONSIDERATIONS (IF APPLICABLE)

You will receive an email when sites complete local considerations. Go to the **Status Summary** page to review.

- Pre-screen** the surveys for completion by clicking the 3/3 Surveys Complete dropdown to view the survey list, then click on the HRP Survey and PI Survey.
- Verify** consent forms are uploaded to the PI Survey, if applicable, and that they are correct. If changes or clarifications are needed, the PI and Study Team Member (or HRPP Liaison) at the site can edit the PI Survey. Only the HRPP can edit the HRP survey.
- Click Export Data** and select **Export Local Considerations** to download a zip file of sites' completed local considerations. Select the site(s) you need and save their files.
- Submit** the site's files to the sIRB for review. Contact the sIRB about submission requirements.

Site	Agreements	Reliance Decision	Local Considerations	Approval Status
Anderson Medical Center	2 / 3 Agreements Complete	Notify & Grant Access		
Central Ohio Medical Center	3 / 3 Agreements Complete	Notify & Grant Access		
Faraday Institute of Research, Science and Technology	3 / 3 Agreements Complete	Contacted 4/13/2023		
Mellon University Medical Center	3 / 3 Agreements Complete	Completed 4/21/2023	3 / 3 Surveys Complete	Not Approved
Middle-earth College of Agriculture	3 / 3 Agreements Complete	Started 4/21/2023		Approved
University of the Bay	3 / 3 Agreements Complete	Completed 5/16/2023		Approved

## STEP

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## UPLOAD INITIAL SIRB APPROVALS FOR PARTICIPATING SITES

- Click **Upload Relying Site Approval** on your Getting Started Checklist or click the **Not Approved** status for the site you wish to upload approval for on the **Status Summary** page.
- Select the site** from the list of sites on the left panel.
- Change** the **Status** to 'approved' and required items will turn red.
- Indicate** the **Review Type** (Full Board or Expedited).
- Is Site Enrolling?** Defaults to **Yes**, change to **No** if the site will not be enrolling participants and does not need a consent form.
- Enter** the **Date Submitted**, **Reviewed**, and **Approved**
- Upload** the site's **IRB Approval Documentation**, **Consents & Assents** (or waive), and other site-specific IRB approved documents.
- Click **Save**.

The site HRPP and study team are notified by email when new approvals are saved, and you are cc'd.

**Relying Site Approvals**

**Vanderbilt Univ**

**Status**: approved

**Review Type**: Initial Study: Full Board

When you save this approval, an email will be sent to the relying site's HRPP and study teams. If you are not ready to notify the site of their approval, change the Status to pending.

**Is Site Enrolling?**  
☒ Yes ☐ No

**Date Submitted**: mm/dd/yyyy (Required)

**Date Pre-Reviewed**: mm/dd/yyyy

**Date Reviewed**: mm/dd/yyyy (Required)

**Date Approved**: mm/dd/yyyy (Required)

**Site Specific Documents**

**IRB Approval Documentation**: Choose a file or drag it here. (Required)

**Consents & Assents**: Choose a file or drag it here. (Required)

## ADDITIONAL RESOURCES

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A. **Uploading other approvals** (see quick guides below)

- [Continuing Review](#)
- [Study-wide Amendments](#)
- [Site Amendments](#)

B. **Site closures**

- Closing a site ensures that only active sites retain access to ongoing studies.
- [Site Closure Quick Guide](#)

C. **Study closures**

- Closing a study ensures all sites are aware that the study ended but retains a record of the reliance and a history of sIRB site approvals.
- [Study Closure Quick Guide](#)